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Introduction

Breast cancer is one of the leading causes of death among women. However, there is clear evidence that early diagnosis and subsequent treatment can significantly improve the chance of survival for patients with breast cancer.¹⁻⁴

Mammography has become one of the major diagnostic procedures with a proven capability for detecting early-stage, clinically occult breast cancers.⁵⁻⁸ However, breast cancers in their early stage are small and frequently their radiographic appearance differs only subtly from that of normal tissue or benign abnormalities. Because of this subtlety, the potential for misclassification by radiologists is substantial. Only 10-30% of cases that have mammographically suspicious findings and are subjected to biopsy prove to be malignant.⁹ On the other hand, approximately 10-30% of patients with breast cancer are misdiagnosed by mammography (have the cancer missed or not detected on their mammograms).¹⁰⁻¹⁴

Besides the subtle nature of radiographic lesions associated with breast cancer, many errors in radiological diagnoses can be attributed to human factors such as subjective or varying decision criteria, distraction by other image features, and simple oversight.¹⁵⁻¹⁷ Studies suggest that these errors may occur even with experienced radiologists.^{18, 19} These errors may be reduced by the use of automated detection schemes that can locate and classify possible lesions, thereby alerting the radiologist to examine these areas with particular caution. Moreover, the automated detection schemes can serve as a "second radiologist", similar to the double reading by two radiologists that is commonly practiced in diagnostic radiology to increase diagnostic efficacy.

Microcalcifications are commonly considered to be important signs of breast cancer. It has been reported that 30-50% of breast cancers detected radiographically demonstrate microcalcifications on mammograms.²⁰⁻²⁵ Up to 90% of cases of ductal carcinoma in situ present with microcalcifications.²⁶ The correlation between the presence of microcalcifications and the presence of breast cancer suggests that accurate detection of microcalcifications will improve the efficacy of mammography as a diagnostic procedure.

Microcalcifications occur in malignant and benign conditions. Some microcalcifications are characteristically benign or are associated with a benign process. For example, calcified fibroadenomas have a typical "popcorn" configuration appearing coarse and solitary. Milk of calcium demonstrates sedimentation.²⁷ Vascular calcifications have a tram track appearance, typical of vascular calcifications seen in other areas of the body. Dermal calcifications tend to be smooth and round with lucent centers. Secretory calcifications are thick, smooth, cigar-shaped, and usually non-branching. Features supporting benignity include uniform size and density of the calcium flecks, as is seen in sclerosing adenosis.²⁸ Furthermore, benign microcalcifications tend to be uniformly dense or scattered, without a segmental or linear distribution.²⁹

Some microcalcifications associated with malignancy have a typically granular or linear appearance. They usually occur in clusters consisting of greater than 15 particles.³⁰ The particle size is small (less than 1 mm) and the shape is irregular.³¹ Some clusters of microcalcifications have neither the typically benign nor typically malignant configurations

described above. These "indeterminate" microcalcifications present a significant diagnostic problem and require careful analysis.

The number of microcalcifications per cm^2 has been shown to be the most important predictor of malignancy, with clusters consisting of less than 10 microcalcifications per cm^2 having a high chance of benignity. Clusters consisting of microcalcifications numbering greater than 15 per cm^2 have a higher chance of malignancy.³²

The task of detection and classification of microcalcifications for the diagnosis of breast cancer is a difficult one. The inability to correctly predict cancer is not only due to the overlap in appearance between microcalcifications associated with benign and malignant conditions, dense breasts, improper technical factors or simple oversight by radiologists may contribute to the failure to detect microcalcifications. Differing levels of confidence and training among interpreting radiologists may lead to inconsistent recommendations for management.

Radiologists classify breast microcalcifications into one of three groups: benign, likely malignant, and indeterminate. Most patients with indeterminate types of calcifications undergo a breast biopsy to exclude cancer. Any method that would correctly classify benign types of calcifications previously considered indeterminate would decrease the frequency of biopsy and therefore the cost of detection of breast cancer.

Several investigators have been developing computer programs for the automated detection of microcalcifications on mammograms.³³⁻³⁶ Chan *et al.* showed that the computer program can detect subtle microcalcifications that may be missed by radiologists, indicating that it is a promising approach to the automated detection of microcalcifications.³⁷ More recently, Wu *et al.* applied an artificial neural network (ANN) to detect microcalcifications.³⁸ The ANN, trained by using the power spectrum of regions of interests (ROI) containing microcalcifications, was able to eliminate 50% of false-positive detections of a rule-based scheme^{37, 39} while preserving more than 95% of the true-positive detections. The neural network achieved an A_z value of 0.85 for the detection of clustered microcalcifications. Several other computer schemes for detection of microcalcifications were also reported by Astley *et al.*,⁴⁰ based on likelihood estimators, by Grimaud *et al.*,⁴¹ using mathematical morphology tools, and by Karssemeijer,⁴² using a stochastic method based on Bayesian decision theory.

As stated earlier, microcalcifications can be associated with either benign or malignant processes. It is important to distinguish different types of microcalcifications after they have been identified by a detection scheme. Accurate classification of microcalcifications into benign and malignant groups would help improve the sensitivity of the diagnosis as well as reduce the number of unnecessary biopsies.

As the first step in the process of developing an automated computer scheme for classification of microcalcifications, a neural network system was developed to classify microcalcifications in the radiographs of biopsy specimens. Classification of microcalcifications in radiographs of biopsy specimens is an "idealized" situation.

In biopsy specimens, underlying tissue around microcalcifications is less than that present in normal mammograms. Therefore, the scatter radiation recorded on films is reduced, resulting in improved contrast. Higher dose and geometrical magnification can also be used to obtain radiographs of biopsy specimens as compared with regular mammograms. Less magnification results in less geometrical unsharpness. Higher exposure can be used to achieve greater signal to noise ratio and thereby improve image quality of radiographs.

Therefore, microcalcifications in the radiographs of biopsy specimens are more clearly represented than those in regular mammograms. After we can successfully apply our algorithm to classify microcalcifications in radiographs of specimens, we will make necessary adjustments to apply the algorithm to the regular mammograms.

In recent years, rapid progress of research on artificial neural networks (ANN)⁴³ has been reported extensively in the field of computer science and many applied fields. Neural networks address detection, classification, and decision-making problems not by pre-specified "conventional" algorithms, but rather by "learning" from examples presented repeatedly. The popularity of neural networks is primarily due to their apparent ability to make decisions and draw conclusions when presented with complex, noisy, or partial information and to adapt their behavior to the properties of the training data. Neural networks are capable of parallel-processing a large amount of information simultaneously and have been shown^{44, 45} to be a useful tool for pattern recognition in fields where conventional algorithmic approaches and rule-based expert systems may not be successful.

Conventional diagnosis of breast cancer, a combination of mammography and physical examination has been the most effective screening methods to date. However, the ability of the conventional techniques to discover cancer with an acceptable reliability is limited. The limitations are manifest in the facts that average positive predicative value for suspicious breast lesions is only 20-30% at surgical pathology and approximately 10-30% of breast cancer are missed by mammography.^{10-14, 46} In addition, mammography may fail to reveal breast lesions that are present in certain groups of women such as those with mammographically dense breasts, women with prior breast cancer who are at risk for recurrence following therapy, and those with prosthetic breast implants.

Recent studies have suggested that BMRI may be an effective tool that complements the conventional techniques such as mammography in the diagnosis and management of breast carcinoma.⁴⁷⁻⁵⁰ BMRI has a very high sensitivity in detecting cancer lesions. The applicability of MRI as a diagnostic procedure in breast cancer is currently limited, however, by the low specificity and high cost. While BMRI shows promise in solving some of the difficult diagnostic problems, the time required to examine the massive amounts of data resulting from each patient can potentially make cost of MRI very high. Technical advances have decreased the data acquisition time to approximately fifteen minutes per patient. However, data analysis remains cumbersome at best and a stumbling block to ready clinical acceptance. It can take 1-2 hours for a radiologist to analyze the large amount of images of each patient. The analysis performed often have qualitative and subjective properties, which is one of the reasons for the low specificity of BMRI.

We have developed an image display and analysis system that allows interactive 3D image manipulation and qualitative analysis of selected image regions. This computer visualization system can help radiologists improve the efficacy of examining the massive amount of data. The computer system may improve the specificity of detecting breast carcinoma, reduce the subjectivity of radiologists in interpreting BMRI images, and cut the cost by reducing the time radiologists have to spend reading the large number of images. This system can make BMRI a cost-effect procedure with high sensitivity and specificity in the diagnosis of breast cancer.

Body of Report

METHODS

Algorithms for classification of microcalcifications have been developed in two stages. First, a convolution neural network based method was developed to classify microcalcifications using radiographs of pathological specimen. Second, an image feature based method was developed to classify microcalcifications using real mammograms. The image features used and optimal structure of the backpropagation neural network were determined by using genetic algorithm.

Classification of microcalcifications in pathological specimen

The overall approach for the classification of microcalcifications using a CNN system is shown in Figure 1. The radiographs of pathological specimen are digitized by a high resolution digitizer. Regions of interest (ROI) containing microcalcifications are manually selected. These ROIs are preprocessed and used as input to the CNN system. Finally, the classification results are examined by the ROC analysis.

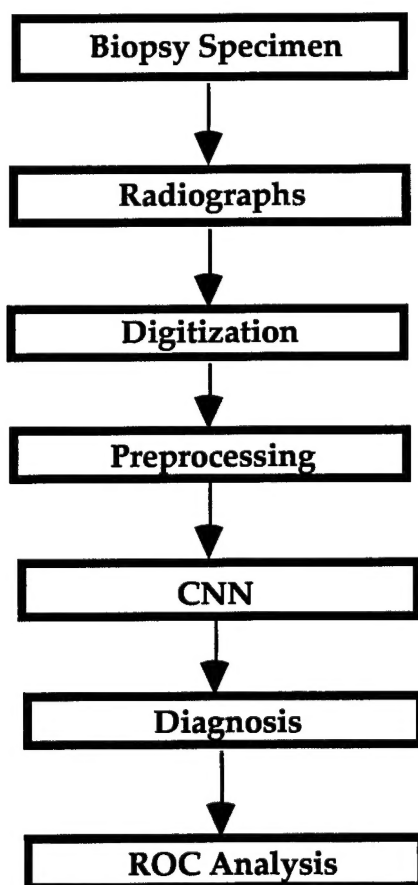


Figure 1. Overall approach for the classification of microcalcifications using CNN.

The classification of microcalcifications is based primarily on the fact that microcalcifications associated with malignant processes generally have more irregular shapes with fuzzy and spiculated boundaries and are less uniform in density and size. They are usually grouped into multi-particle clusters. The microcalcifications associated with benign processes, on the other hand, usually tend to have smoother and well-defined boundaries, rounded shape, and uniform densities and sizes. The neural network system will be trained to recognize the characteristics of each type of microcalcifications.

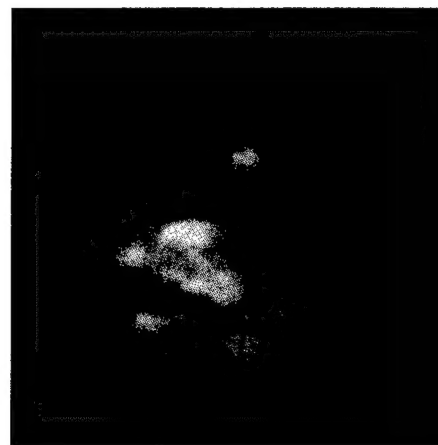
The CNN is based on the network structure of Fukushima's Neocognitron⁵¹ which is designed to simulate the vision of vertebrate animals. The structure of CNN used in this study resembles a simplified Neocognitron. A two-dimensional convolution operation from the input layer to the hidden layer is employed to simulate radiologists' viewing of a suspected area. The CNN has the ability to process and recognize two dimensional image patterns and has been shown to be an effective tool in image processing and pattern recognition.⁵²⁻⁵⁴

Acquisition of Mammograms

The selected radiographs of breast biopsy specimen are digitized with an image resolution of $21\mu\text{m} \times 21\mu\text{m}$ per pixel by a CCD camera digitizer (DBA Systems Inc.). With high resolution digitization, the morphological information of microcalcifications can be preserved, which enables the neural network system to differentiate different types of microcalcifications on the basis of their geometrical shapes and density patterns. Figure 2(a) shows a cluster of microcalcifications in original radiographs of pathological specimen. Shown in Figure 2(b) are clustered microcalcifications after being digitized with the high resolution digitizer. The shapes and density patterns of the microcalcifications are better defined in Fig. 2(b) than those shown in Fig. 2(a).



(a)



(b)

Figure 2. Microcalcifications shown in an original radiograph of pathological specimen (a) and shown in a radiograph digitized with a high resolution ($21\mu\text{m} \times 21\mu\text{m}$) digitizer (b).

Convolution Neural Network

The structure of the CNN⁵² is shown in Figure 3. The input to the CNN are ROIs of matrix size of 64×64 pixels, containing benign or malignant type of microcalcifications. Only one hidden layer is used in this study. The connections between input and hidden layer are grouped into seven different kernels based on the structure of Fukushima's Neocognitron.^{51, 55} There are two output units in the output layer, with each unit corresponding to a benign or malignant class of microcalcifications. The hidden layer and the output layers are fully connected.

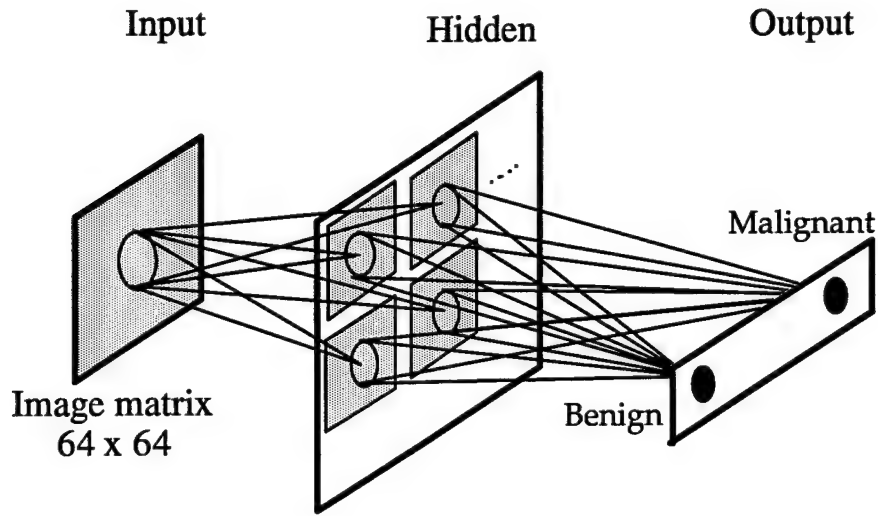


Figure 3. Structure of a convolution neural network used for the classification of microcalcifications.

The training algorithm of the CNN is similar to that of a backpropagation neural network, in which backpropagation and the generalized delta rule are used in the training process.⁵⁶ The input signals are now two dimensional images. The weights are all arranged in the convolution kernels. In the feed-forward propagation, the output of l th layer are first convoluted with weight filters. The sum of the convolution is then added by a bias term to form the net input to the next layer.

$$N_q^{l+1}(x,y) = \sum_{p=1}^{P^l} [O_p^l * W_{p,q}^l]_{(x,y)} + b_q^{l+1} \quad [1]$$

where $N_q^{l+1}(x,y)$ is the net input to the unit (x,y) in layer $l+1$, $O_p^l(x,y)$ is the output of the unit in layer l , $W_{p,q}^l(x,y)$ is a weight kernel, and b_q^{l+1} is a bias term in layer $l+1$. In the notation, layer number $l = (1, 2, \dots, L)$, cluster number in the l th layer $p = (1, 2, \dots, P^l)$, and cluster number in the $(l+1)$ th layer $q = (1, 2, \dots, P^{l+1})$. Note that $*$ denotes discrete convolution,

$$[O_p^l * W_{p,q}^l]_{(x,y)} = \sum_i \sum_j (O_p^l(i,j) \cdot W_{p,q}^l(i-x, j-y)). \quad [2]$$

We can then rewrite equation [1] as

$$O_q^{l+1}(x,y) = f(N_q^{l+1}(x,y)), \quad [3]$$

where f is the activation function

$$f(x) = \frac{1}{1 + \exp(-x)}. \quad [4]$$

In the error backpropagation, the weights are modified, similar to that in Eqn.[2], as the following,

$$\Delta W_{p,q}^l(n+1) = \eta(d_q^{l+1} * O_p^l) + \alpha \Delta W_{p,q}^l(n), \quad [5]$$

to minimize the error function,

$$E = \frac{1}{2} \sum (T(x,y) - O_1^L(x,y))^2, \quad [6]$$

where $T(x,y)$ is the target output.

In the training process of the CNN, each image block is rotated and reflected such that the number of training data are increased eight fold. The rotation and reflection represent different orientations of microcalcifications in mammograms. The training with additional orientation can effectively make the CNN rotational invariant.

Classification of microcalcifications in real mammograms

An alternative approach was employed to classify microcalcifications of in clinical mammograms. Microcalcifications can be characterized by a number of quantitative variables describing the underlying key features of a suspicious region such as the size, shape, and number of microcalcifications in a cluster. These features are calculated by an automated extraction scheme for each of the selected regions. The features are then used as input to a backpropagation neural network to make a decision regarding the probability of malignancy of a selected region. The initial selection of image features set was a rough estimation that may include redundant and non-discriminant features. A genetic algorithm was employed to select an optimal image feature set from the initial feature set and select an optimized structure of the neural network for the optimal input features. Finally, the performance of optimized neural network is evaluated using the selected sub-features. The performance of neural network is then compared with that of radiologists in classifying the clusters of microcalcifications.

A selected region is a 256×256 matrix from a mammogram that is digitized at 50 microns per pixel with a Lumisys scanner. Figure 4 shows two regions of interest (ROI) selected from original mammograms: (a) benign and (b) malignant. The background trend in a selected region is eliminated by an adaptive trend correction technique. The trend-corrected image is then binarized by taking the top level of 32 quantization levels. A series of pre-processing techniques that include open and dilation operations, logic AND, and labeling operation are applied to the binary image to eliminate artifacts while preserving the original shapes of microcalcifications. Image features are extracted based on the pre-processed image and two morphologically filtered images. Nineteen image features are extracted for each cluster of microcalcifications characterizing the size, number of microcalcification in the cluster, and shape.

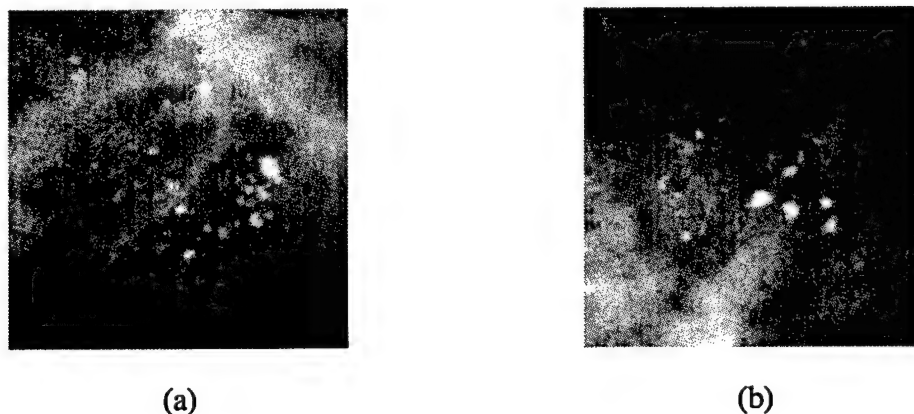


Figure 4 Two ROIs of size 256×256 selected from original mammograms: (a) benign and (b) malignant

Pre-Processing

The pre-processing algorithm is demonstrated in Figure 5. The purpose of pre-processing is to obtain a binary image of microcalcifications. The pixels representing microcalcifications have values of one and the rest have values of zero. We need only binary image because image features we use are not based on density but rather based on only size, shape, clustering, and number of microcalcifications in a cluster.

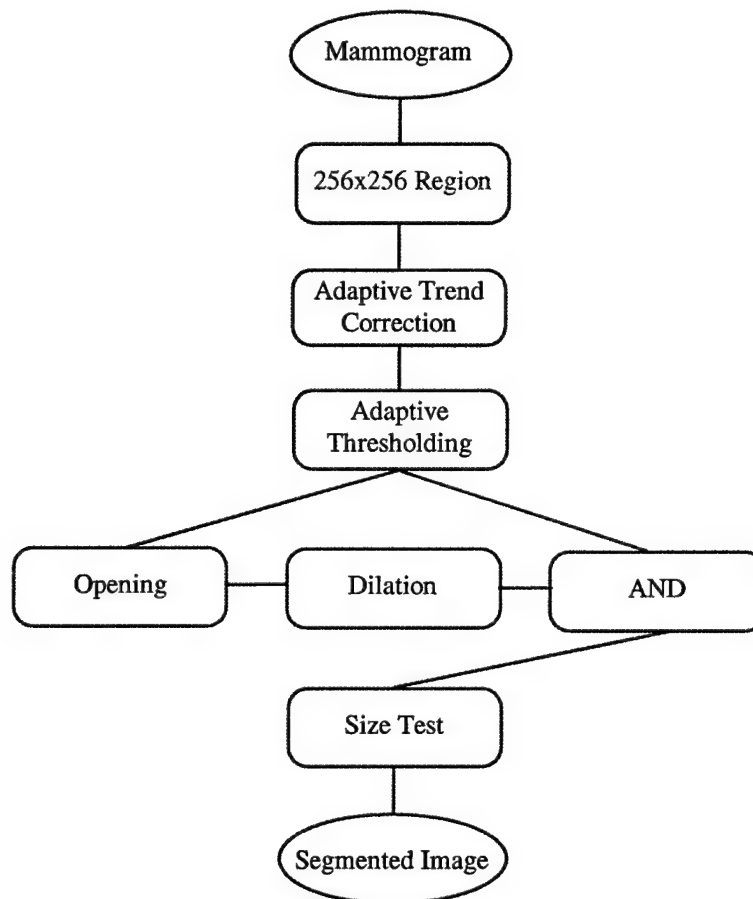


Figure 5 Pre-Processing of Selected Region of a Mammogram

a. Adaptive background trend correction

A average filter of kernel size of 23×23 pixels is used to remove the background trend in a selected region. At each pixel in the selected region, an average value is calculated for the surrounding 23×23 pixels and subtracted from the original pixel value. This process is repeated for all the pixels in the selected region. The adaptive technique is used to isolate microcalcifications from their immediate surroundings rather than from a different region that may have a different density level. As long as the sizes of microcalcifications remain small and microcalcifications are not clustered too close to each other, this method will be able to preserve the original shapes of microcalcifications. The kernel size of 23 pixels is equivalent of approximately 1.2mm. Therefore, microcalcifications that are larger than 1.2mm in size are eliminated by the background correction process.

b. Adaptive thresholding

After background correction, a threshold is applied to the image such that only 2048 pixels of highest intensity (pixel value) are kept (assigned pixel value 1) and the rest pixels are assigned pixel values of 0. The number of pixels that are kept after thresholding is determined empirically that the sizes of microcalcifications in the threshold image are similar to those in the original image.

c. Opening operation

A morphological Opening operation with 3×3 kernel size is applied to the binary image to remove line artifact. The Opening is a combination of an erosion followed by a dilation.

d. Dilation

A morphological Dilation operation is applied after the OPEN operation to fill in the small holes near the boundary of an object.

e. AND operation

To preserve the shape of microcalcifications, an AND operation between the original binary image and the image after dilation is performed.

f. Labeling

A labeling process is applied after the AND operation to eliminate small objects that still remain in the image that are less than 5 pixels in size. Any object that is that small is unlikely to be an microcalcification.

Image Feature Extraction

Three images are used in feature extraction. The three images are: Image1, the preprocessed image; Image2, dilation of Image1 by a 5×5 kernel; Image3, dilation of Image1 by a 25×25 kernel. The process is demonstrated in Figure 6.

1. Number of microcalcifications N from Image1
2. Area (number of pixels) in Image1 $S1$
3. Area (number of pixels) in Image2 $S2$
4. Area (number of pixels) in Image3 $S3$
5. $(S2-S1)/N$, a measure of average number of irregular pixels
6. $(S3-S1)/N$, a measure of average distance between microcalcifications in a cluster
7. $(S2-S1)$, total number of irregular pixels

8. (S3-S1), a measure of average distance between microcalcifications in a cluster
9. V1, size of the largest microcalcification in Image1 - size of average microcalcification in Image1
10. V2, size of the average microcalcification in Image1 - size of the smallest microcalcification in Image1
11. V3, size of the largest microcalcification in Image1
12. V4, size of average microcalcification in Image1
13. V5, size of smallest microcalcification in Image1
14. E1, ellipticity of the largest microcalcification on Image1
15. WAE, weighted average ellipticity of microcalcifications on Image1
16. A_x , length of semi-major axes of the fitted ellipse of a cluster on Image1
17. B_x , length of semi-minor axes of the fitted ellipse of a cluster on Image1
18. A_x / B_x
19. $A_x \times B_x$

A total of 19 image features are extracted for candidate cluster and used as input to a backpropagation neural network.

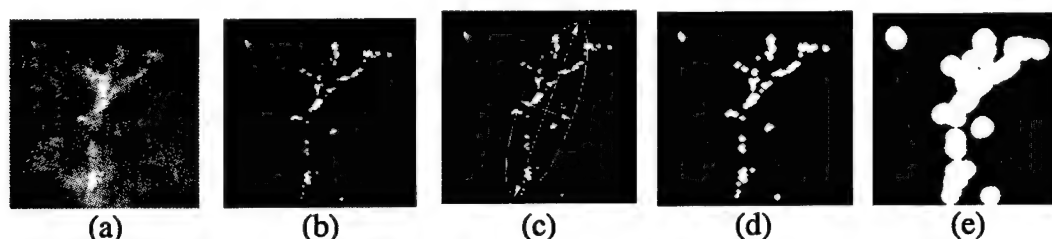


Figure 6 (a) Original gray scale image, (b) Binary image, (c) Best-fit ellipse and its semi-major axis A_x and semi-minor axis B_x , (d) Dilation with 5×5 kernel, and (e) Dilation with 25×25 kernel.

Genetic Algorithms

Genetic algorithms are directly modeled after biological systems and behavior. All the values of a structure represent the neural network characteristics that uniquely define a candidate solution in the space of possible solutions. Utilizing genetic algorithms, the Genetic Supervisor evolves successor populations, or generations, from a limited population of initial candidate solutions. It does this by treating the inclusion or exclusion of each column of data

in the full Input column set as features; the number of layers and the number of neurons per layers as features; and the control parameters of each neural network as features.

These features are then varied in each new generation with the resulting structure evaluated in terms of neural network fitness. Each structure in the generation is evaluated and judged by either the lowest RMS Error achieved after a fixed number of epochs or by the number of epochs taken to achieve a minimum point in RMNS Error. These two measures represent neural networks that train to minimal error or neural networks which train with minimal epochs. These criteria can be applied to the set of training cases or the set of test cases.

If the structure representing a neural network successfully meets the fitness criteria selected, then the values of its features will be retained and bred with other structures. For each generation, the Genetic Supervisor generates a population of structures in one of two ways. All the structures of an initial generation and a certain number of structures in subsequent generations are created with features set to random values constrained within specified limits. Subsequent generations are created by cross-breeding the strings of successful structures or occasional mutations of randomly selected features of successful structures. Some or many of the weakest structure may be culled, these are replaced with new structures.

Through this evolution-like process, an optimal neural network can be developed. Note, however, that this process requires the training of many versions of the neural network to determine an optimal one; for neural network models that have large network configurations or have large data sets this can be a lengthy process – but so can biological evolution!

A system consisting of a Backpropagation neural network and a Genetic Supervisor is developed to classify the extracted image features. The Genetic Supervisor attempts to select the best subset of image features from the provided input set, configure the best neural network structure, and adjust the parameters of the network for optimum performance. The performance of the neural network system is evaluated by ROC analysis.

Reduction of Patient Call Backs

A third set of images will be selected at GUMC which are images of patients that are called back after screening exam to follow up. A majority of the called back patients are benign. The objective is to reduce the number of call back patients are definitely benign and therefore reduce the overall cost of breast cancer diagnosis practice.

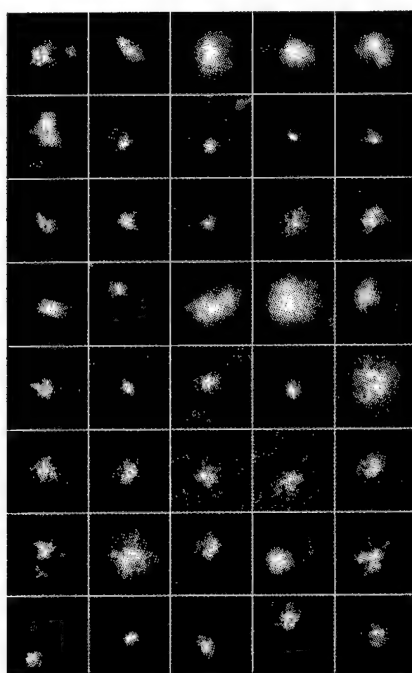
Database

Pathological Specimen Images

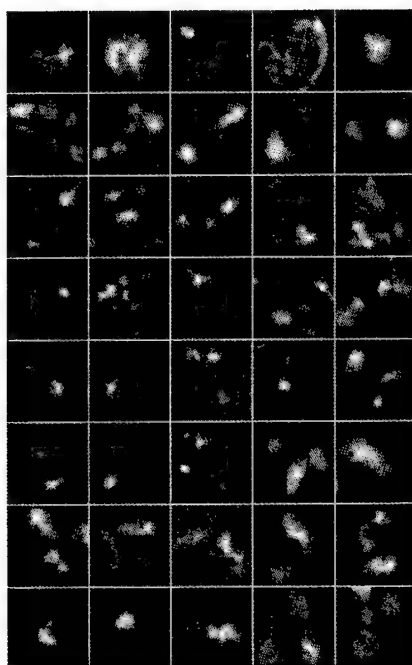
Eighty regions of interest (ROI) that contain clustered microcalcifications (40 benign and 40 malignant) are selected from 60 digitized radiographs of pathological specimen in this study. Figure 7 shows all of the 80 ROIs selected in the database. There are substantial variations in size among benign or malignant microcalcifications. The information concerning the classification of microcalcifications ("truth") are obtained from the results of biopsy examination. Background trend correction is employed to remove the non-uniform background structure in different ROIs.

Clinical Mammograms

Two sets of clinical mammograms were collected. The first set contains 70 cases of clusters of microcalcifications and the second set contains 62 cases of microcalcifications. The two sets cases were selected from two different sources of medical institutions. The microcalcifications in both sets were rated subtle by radiologists. The diagnostic truth of all of the cases in the selected sets had been verified by biopsy.



(a)



(b)

Figure 7 Database for the training and testing of the CNN; (a) 40 ROIs containing benign clustered microcalcifications and (b) 40 ROIs containing malignant clustered microcalcifications.

Diagnosis of breast cancer by MRI using a neural network based computer vision system

Figure 8 shows the overall scheme of the two-step approach that is employed in this project. After the images are acquired, an interactive image display and image analysis is used to aid the radiologists' examination of the images and initial diagnoses. Meanwhile, the computer vision system will detect the suspicious areas in the 2D slices. The detected lesions at various slice images will then be grouped automatically to form 3D lesions. A 3D convolution neural network will be employed to distinguish the 3D lesions that have shapes typical of malignant tumors from other benign lesions. The final results will be presented to radiologists to make a final diagnosis.

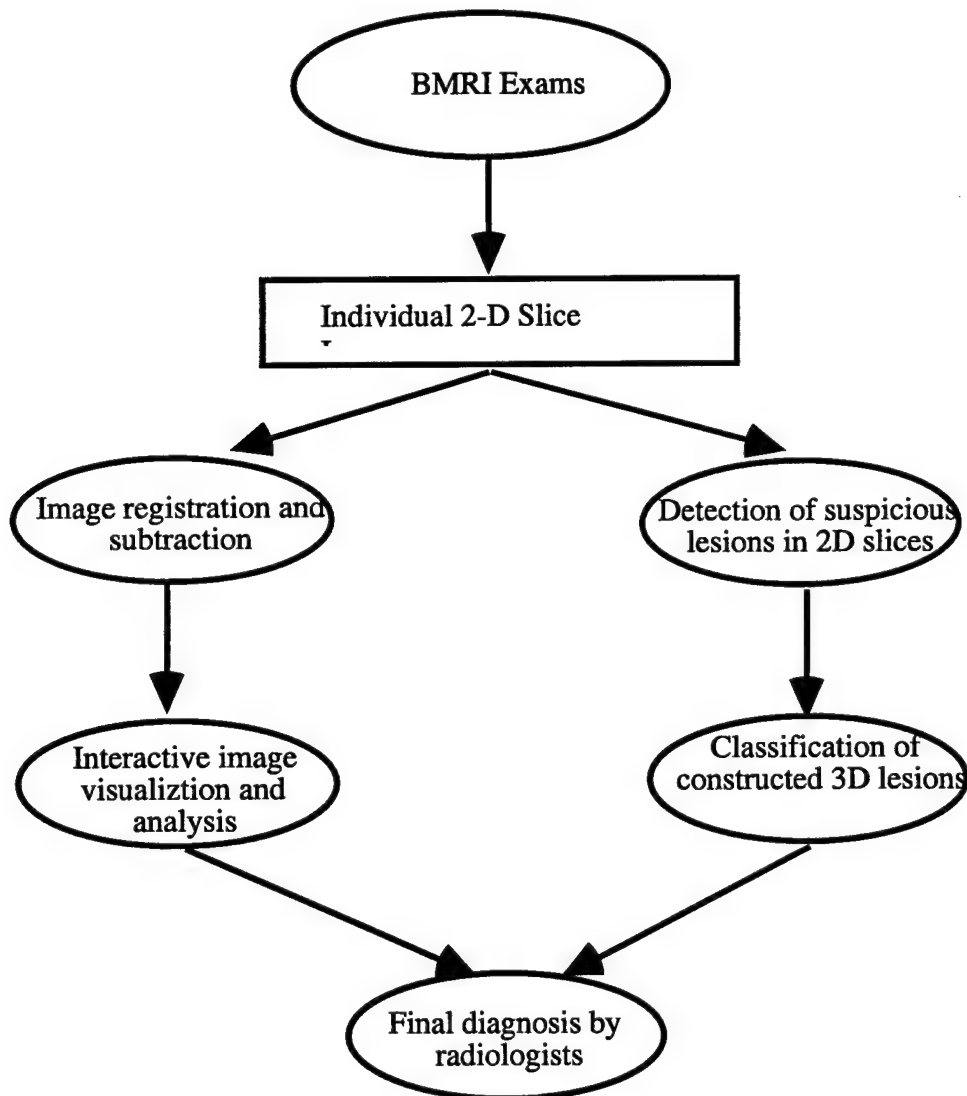


Figure 8 Overall scheme of the neural network based computer vision system

Image acquisition and display

The BMR examinations are performed using a bilateral dedicated breast coil operating in the receive mode with a 1.5 Tesla MRI system. The imaging pulse sequence used is a 3D FISP (fast imaging with steady state free precession) variant, a rapid gradient echo (GRE) technique that allows volume imaging of both breasts with 2.5mm consecution slice thickness. Each acquisition sequence uses 64 partitions for the total slab thickness of 160mm. The matrix size of 2D slice field of view is 256 x 256. The width of the field is approximately 380mm, making the pixel size in the 2D slice image about 1.5mm. The imaging time per sequence is 84 seconds.

The scans are performed once prior to contrast administration and five consecutive times following intravenous gadolinium bolus administration. No delays occur between each of the five contrast-enhanced scans. The images are calculated after all five enhanced scans are completed. The gadolinium contrast agent is administered at a dose of 0.1 mmol/kg using one of the three FDA-approved paramagnetic agents: gadolinium diethylene triamene pentaacetic acid demeglumine (Gd-DTPA, Berlex); gadolinium diethylene triamene pentaacetic acid bis-methyl amide (Gd-DTPABMA, Winthrop); or gadolinium tris (hydroxymethyl) aminomethane (Gd-DO3A, Squibb). The paramagnetic agents are given via the largest indwelling catheter cable to be placed in a peripheral vein over five seconds. The contrast injection will be followed by 10cc saline flush injection to clear the intravenous line of contrast.

We have developed an image registration technique to match image slices at same location during a time sequence so that the pre-contrast image can be subtracted correctly from the post-contrast images. Image slices are stacked up to construct a 3D image object by volume rendering. The 3D object can be rotated and viewed from different angles. The user can also select an arbitrary projection plane and display the density of breasts in the selected plane.

3D images in one complete exam before and after contrast injection can be displayed in a continuous loop to demonstrate the dynamic change of signal enhancement in the lesions. The 3D display of subtracted images after contrast injection will reveal regions of strong signal enhancement. Radiologists can then zoom in to those regions of strong enhancement and analyze them with quantitative image analysis tools that will be developed in this system. The image analysis tools can plot enhancement curves at any given location.

Detection of suspicious areas in images of 2D slice images

Enhancement profiles that show the signal intensities at different time intervals before and after the injection of contrast agent will be calculated at each pixel in a slice image. Features characterizing the signal enhancement after contrast injection will be automatically extracted from the enhancement profiles. The features will be calculated based on time derivatives of the enhancement curve. The feature selection will be optimized by using a self-organizing feature map neural network.

A three-layer, feed-forward, back-propagation neural network will then be employed to recognize pixels with abnormal enhancement profiles. Enhancement profiles will be selected from cancer and normal regions of the breasts to train the neural network. The trained neural

network, provided with enhancement profiles of individual pixels of an image, will be able to identify those pixels belonging to a region that could be potentially cancerous. A 2D lesion is formed by connecting all the positive pixels identified by the neural network in a local area in a 2D slice image. The detected lesions in consecutive 2D slice images will be grouped together using a region growing technique to construct 3D objects.

Classification of 3-D suspicious lesions

Suspicious regions that are identified in the previous step will include both malignant and benign lesions. Studies have shown that the morphology of lesions may present a clue to lesion origin.⁵⁷ The presence of internal septation and lobulation suggest the lesion represents a fibroadenoma, while border irregularity and rim enhancement are highly suggestive of carcinoma. The morphology of lesions is better delineated in 3D than in single 2D slice images. A three-dimensional convolution neural network (CNN) will be employed to recognize image patterns corresponding to benign and malignant lesions. CNN has been shown to be an effective tool for two dimensional image pattern recognition.⁵⁸ We have applied CNN in many different applications to classify various image patterns in chest x-ray images and mammograms. CNN has been found to be superior to the conventional fully connected neural networks in processing two-dimensional images of relatively large matrix sizes.⁵⁹

The suspicious areas detected in a slice image are two-dimensional projections on a arbitrary plane of three-dimensional lesions. As discussed earlier, the morphology of lesions may be important information in distinguishing benign lesions from malignant cancers. However, spiculated boundaries of three dimensional lesions may not show up in some of their two dimensional projections. As a result, lesions may not be classified correctly based on their projections on a 2D plane alone. To improve the specificity of detecting cancer lesions, suspicious areas that are identified in each of 2D slice images will be grouped together to form 3D lesions. The 3D lesions will be classified into benign and malignant types of lesions by using a three dimensional convolution neural network. The classification will be based on the three-dimensional boundary information of the lesions.

The structure of the 3D neural network is demonstrated in Figure 2, with the convolution (the connections between the input layer and each kernel in the hidden layer) carried out in three dimensional space. The 3D convolution neural network is developed based on the 2D CNN model that has been successfully applied in pattern recognition of digital mammography. In the convolution neural network, the connections between two layers are grouped into a number of clusters, each functioning as a 3D convolution filter. Therefore, the processing of input signals by the CNN is spatially shift-invariant. The shift-invariance makes CNN suitable for image pattern recognition. Pixel values of the 3D objects will be used as input to the CNN.

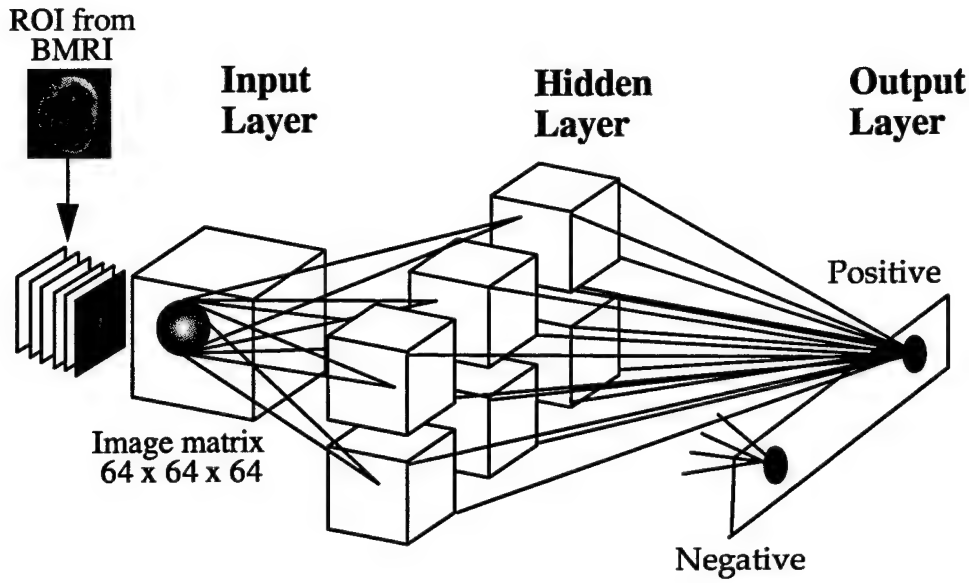


Figure 9 Structure of a 3D convolution neural network

The training algorithm of the 3D CNN^{53, 54} is similar to that of a back-propagation neural network.⁶⁰ The input signals are now three dimensional images. The weights are all arranged in the convolution kernels. In the feed-forward propagation, the outputs of l th layer are first convoluted with weight filters. The sum of the convolution is then added by a bias term to form the net input to the next layer.

$$N_q^{l+1}(x,y,z) = \sum_{p=1}^{P^l} [O_p^l * W_{p,q}^l]_{(x,y,z)} + b_q^{l+1} \quad [7]$$

where $N_p^{l+1}(x,y,z)$ is the net input to the unit (x, y, z) in layer $l+1$, $O_p^l(x,y,z)$ is the output of the unit in layer l , $W_{p,q}^l(x,y,z)$ is a weight kernel, and b_q^{l+1} is a bias term in layer $l+1$. In the notation, layer number $l = (1, 2, \dots, L)$, cluster number in the l th layer $p = (1, 2, \dots, P^l)$, and cluster number in the $(l+1)$ th layer $q = (1, 2, \dots, P^{l+1})$. Note that $*$ denotes discrete convolution,

$$[O_p^l * W_{p,q}^l]_{(x,y,z)} = \sum_i \sum_j \sum_k (O_p^l(i,j,k) \cdot W_{p,q}^l(i-x, j-y, k-z)). \quad [8]$$

In the error back-propagation, the weights are modified, similar to that in a back-propagation neural network,⁶⁰ as the following,

$$\Delta W_{p,q}^l(n+1) = \eta(d_q^{l+1} * O_p^l) + \alpha \Delta W_{p,q}^l(n), \quad [9]$$

to minimize the error function,

$$E = \frac{1}{2} \sum (T(x, y, z) - O_1^L(x, y, z))^2, \quad [10]$$

where $T(x, y, z)$ is the target output.

RESULTS

Images of pathological Specimen

Jackknife Method

A jackknife method was employed to evaluate the performance of the CNN. In the jackknife method, half of the ROIs were randomly selected from the database of 80 ROIs. These ROIs were used to train the convolution neural network. The other half of the ROIs were then used to test the performance of the CNN. By choosing different random samples from the database, the jackknife test can be repeated to generate multiple test output and provide a better estimate of the true performance of the CNN in classifying benign and malignant clusters of microcalcifications.

ROC Analysis

The output values from the two output units were examined by using Receiver Operating Characteristic (ROC) analysis.^{61, 62} The LABROC4 algorithm⁶³ developed by Metz *et al.* was used to fit ROC curves to the continuous data from the output of CNN. The area under the ROC curve (A_z) was used as an overall measure of diagnostic performance. The result from each jackknife test was analyzed individually by using ROC analysis. Ten jackknife tests were performed. A final ROC curve was obtained by averaging the results from the 10 jackknife tests, as shown in Figure 10. The CNN system performed very well in classifying benign and malignant clusters of microcalcifications, achieving an A_z value of 0.90.

Potential Application in Recommending Courses of Action

A potential application of CNN is to classify microcalcifications into groups of definitely benign and possibly malignant. By applying a low threshold level to the output values of the CNN, we can make CNN a classifier that is not very specific but with 100% sensitivity.

With such a classifier, some benign microcalcifications may be classified as possibly malignant, but all of the microcalcifications classified as benign are definitely negative. Thus, radiologists can ignore the clusters of microcalcifications that are classified as benign and only concentrate on those that are classified as possibly malignant. As a result, the time radiologists spend reading mammograms can be reduced and detection efficacy of breast cancer can be expected to improve.

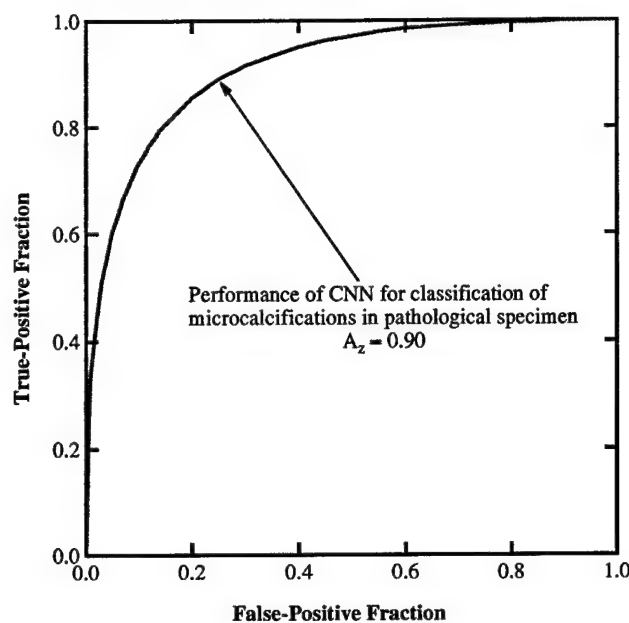


Figure 10 ROC analysis of the performance of CNN in classifying benign and malignant microcalcifications.

The ability of CNN to serve as a classifier to eliminate benign microcalcifications can be demonstrated in Table I. In each of the jackknife tests, we set threshold levels of the output of CNN such that all of the malignant ROIs are to be called positive by the computer system and calculate the number of the benign ROIs that can be called negative (i.e., have output values below the threshold level for positive ROIs).

Table I shows the results for each individual jackknife test as well as the averaged results of the 10 jackknife tests (second column). The CNN can identify, on average, approximately 42% of the benign clusters of microcalcifications with 100% sensitivity (without missing any malignant clusters). Therefore, if this CNN system were used to help radiologists in detecting malignant microcalcifications, radiologists would only need to examine about half of the detected microcalcifications.

As discussed earlier, studies have shown that approximately 10-30% of breast cancers are missed by mammography and only 10-30% of biopsy cases recommended by mammography are actually malignant. As a comparison to the reported performance of radiologists in breast cancer diagnosis, we also listed in Table I (third and fourth columns) the average specificities and positive predictive values, defined as the portion of malignant cases among the biopsied cases.

With a sensitivity of 80%, approximately the same level of sensitivity reported by average radiologists, the neural network system achieved a positive predictive value for malignancy of 84%, compared with 10-30% achieved by radiologists. Therefore, the CNN appears to be a very promising tool for assisting radiologists in making decisions for the diagnosis of breast cancer.

Real mammograms

Genetic algorithms are employed to optimize the structure of a neural network and select the best performing subset from the initial image features. Additional cases of patients that have been recalled for further examination are being collected at GUMC. The performance of the neural network system based on these cases will help determining the usefulness of the system in reducing unnecessary biopsies.

Optimized Feature Set by Genetic Algorithms

Genetic Algorithms were applied to the first subset of seventy cases selected from the database. Two different sets of image features, each consisting of six features, were selected as a result of the optimization process. The neural network performed equally well with the two selected subsets.

Feature Set #1:

1. V3, size of the largest microcalcification in Image1
2. V4, size of average microcalcification in Image1
3. V5, size of smallest microcalcification in Image1
4. E1, ellipticity of the largest microcalcification on Image1
5. WAE, weighted average ellipticity of microcalcifications on Image1
6. A_x, length of semi-major axes of the fitted ellipse of a cluster on Image1

Feature Set #2:

1. E1, ellipticity of the largest microcalcification on Image1
2. WAE, weighted average ellipticity of microcalcifications on Image1
3. A_x, length of semi-major axes of the fitted ellipse of a cluster on Image1
4. B_x, length of semi-minor axes of the fitted ellipse of a cluster on Image1
5. V3 - V4
6. V4 - V5

The optimized parameters for the neural network are: learning rate 0.7 ~ 0.95 and momentum 0.02 ~ 0.06.

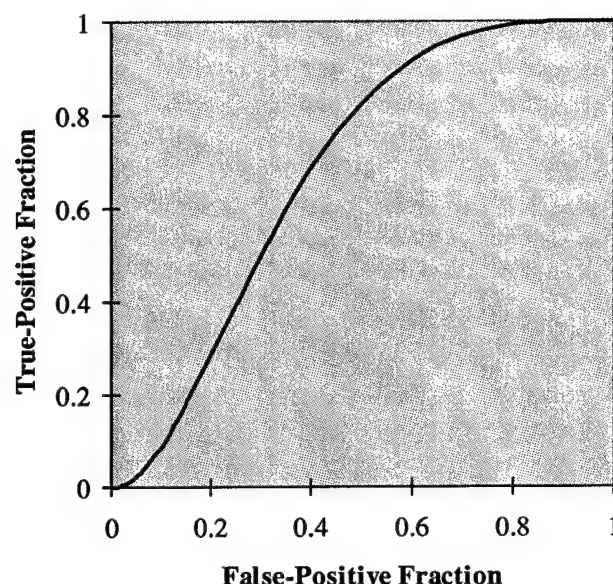


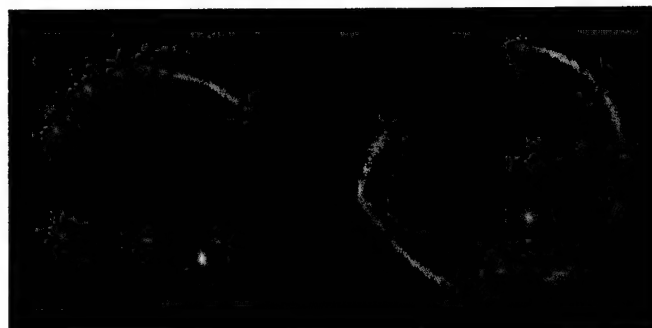
Figure 11 Performance of feature-based neural network in classification of microcalcifications

We evaluated the performance of the optimized neural network by using round-robin method and ROC analysis. Figure 11 shows the ROC curve for the classification of microcalcifications based on extracted image features.

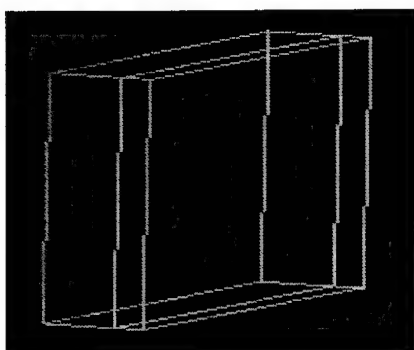
The system of neural network and genetic algorithms improved performance over our previous TRBF neural network⁶⁴. The neural network system was able to classify benign and malignant microcalcifications at a level favorably compared to experienced radiologists. On the same database evaluated, the ROC curve for an experienced radiologist yielded an A_z of 0.54, while the A_z of the ROC curve for the neural network with optimized structure is 0.68. The use of the neural network system can be used to help radiologists reducing the number biopsies in clinical applications.

Case demonstration of 3D image visualization

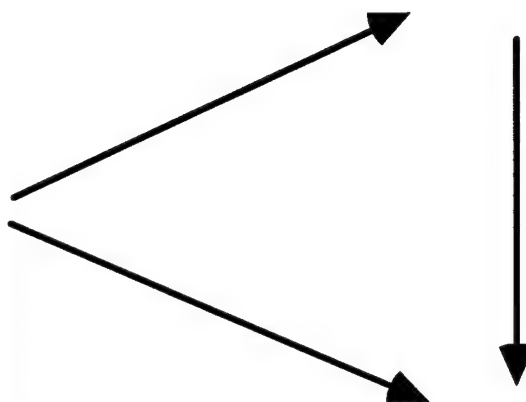
Figure 12 demonstrates a display of selected 2D slice images (a), a constructed 3D image (c), and signal enhancement curve of a selected region (d). The enhancement curves are used by radiologists to determine the probability of malignancy of a specific region. Other characteristics of signal enhancement such as enhance and washout time, gradients of enhancement curves, and variations among a selected region can also be calculated.



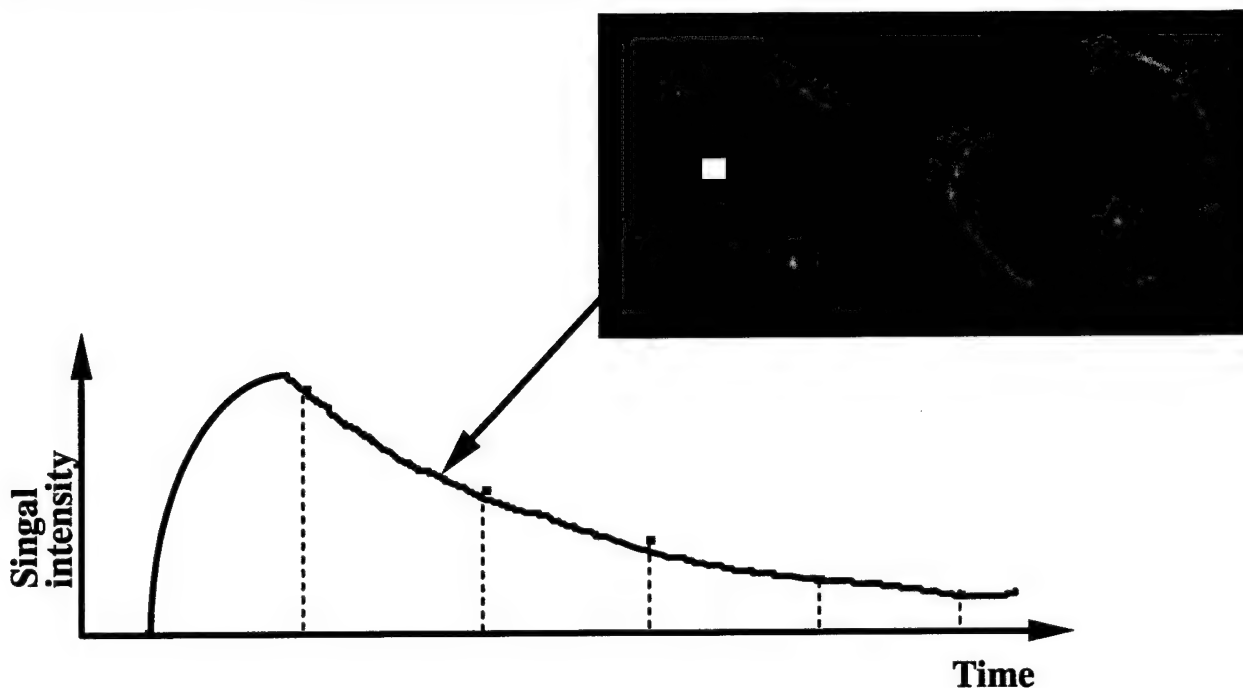
(a) 2D slice image



(b) Position of the 2D slice in the breast



(c) Reconstructed 3D image



(d) Enhancement curve of a selected region

Figure 12 Image visualization in BMRI

Two patient exams are demonstrated here. In each exam, the 64 two-dimensional slices at each time interval are stacked up to construct a three dimensional image. Volume rendering as well as surface-rendering are used to visualize the 3D image object. For each patient exam, there are one pre-contrast 3D image and 5 consecutive post-contrast 3D images. The six reconstructed 3D images are displayed in a continuous loop so that the change of signal enhancement over time can be observed. In each of the 3D images, a 2D projection plane can be selected at any orientation to step through the 3D image and display the signal intensity within the 2D plane. The user can also point to any specific location in a 2D image and display the pixel value at that location. A user can also select a specific region in any 2D plane and display the change of signal intensity over the time sequence to obtain signal enhancement curves.

In the first case demonstrated here, a patient has a tumor in the right breast which is almost invisible in the original images. It can be identified on the subtracted images, as shown in Figure 13.

The second case is a normal case. The image subtraction and 3D reconstruction make it easier for radiologists to locate potential tumor. The radiologists can examine the suspicious area in detail from the 2D projections planes in which the suspicious areas are located. Figure 14 shows the signal enhancement of subtracted 2D slice images.

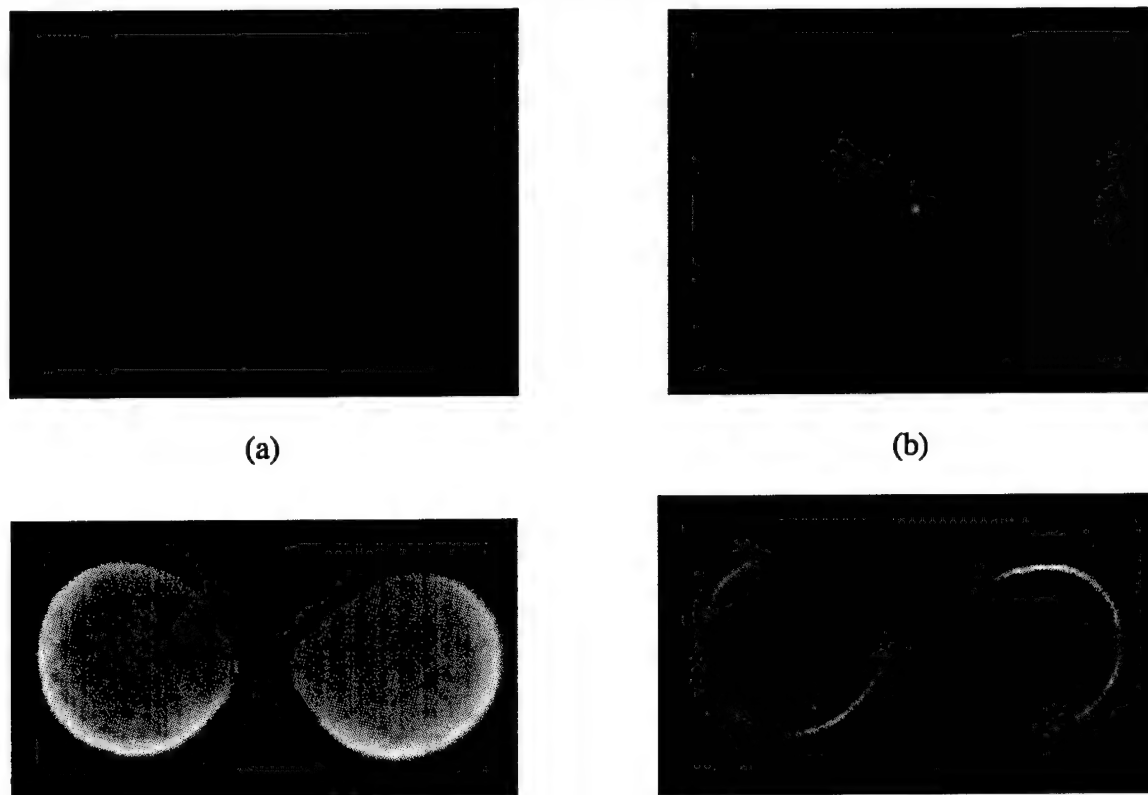


Figure 13 Effect of image subtraction: (a) 3D original image (b) 3D subtracted image (c) 2D original slice (d) 2D subtracted slice

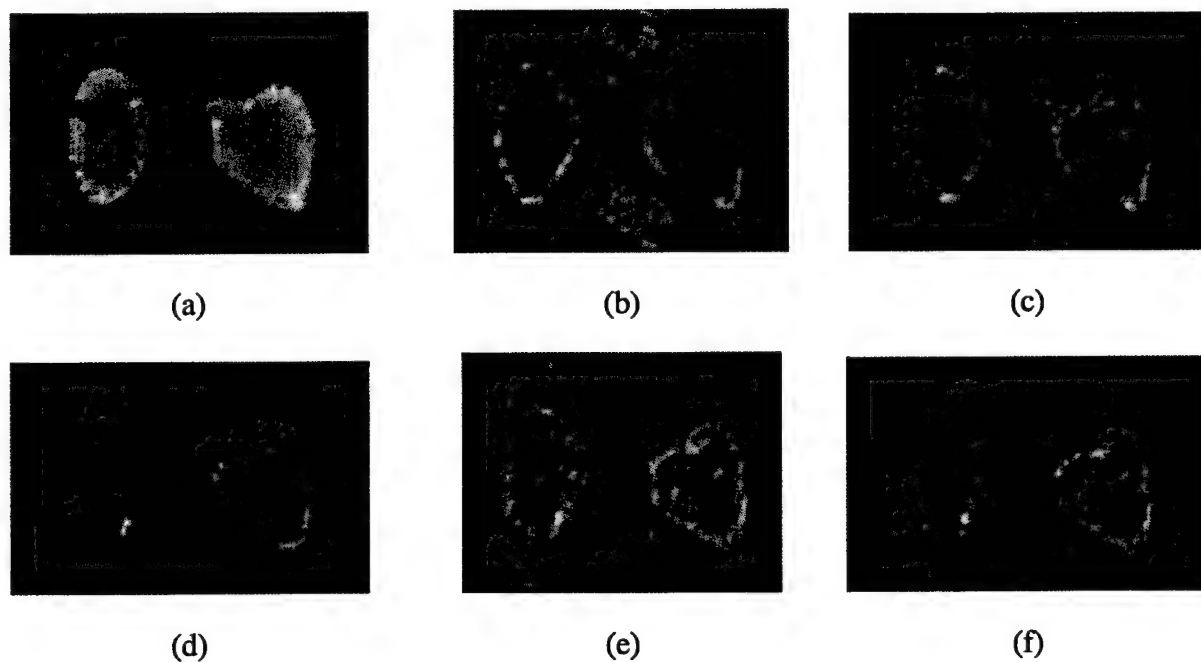


Figure 14 Signal enhancement of breast images in MRI: (a) pre-contrast image (b-f) 5 post-contrast images.

DISCUSSIONS

It is important to note that the CNN is designed as an automated classifier of microcalcifications for the diagnosis of breast cancer. It will be used in conjunction with other schemes for the detection of microcalcifications in digital mammograms. Once microcalcifications are detected, the CNN will be applied to classify them into benign (negative) and malignant (positive) groups. Radiologists can ignore the microcalcifications that are classified into the benign group and examine those that are classified as malignant to decide whether to recommend biopsy or short term follow-up exams.

Table I. Application of CNN in Recommending Courses of Action

| Jackknife Test | Correctly identified negative cases at 100% sensitivity | False Positives at 90% sensitivity | False Positives at 80% sensitivity |
|----------------------------|---|------------------------------------|------------------------------------|
| 1 | 16 | 2 | 2 |
| 2 | 12 | 7 | 2 |
| 3 | 15 | 2 | 0 |
| 4 | 2 | 6 | 6 |
| 5 | 11 | 3 | 1 |
| 6 | 4 | 2 | 2 |
| 7 | 7 | 4 | 2 |
| 8 | 7 | 8 | 8 |
| 9 | 3 | 5 | 5 |
| 10 | 6 | 5 | 2 |
| Average | 8.3 | 4.4 | 3 |
| Average Specificity | 42% | 78% | 85% |
| Positive Predictive Value* | 63% | 80% | 84% |

* *Positive Predictive Value* — Defined as the portion of the actually positive cases among the cases diagnosed that are classified as positive by a diagnostic system.

The results discussed are based on radiographs of biopsy specimen of microcalcifications. The specimen images have, in general, better image quality and greater signal-to-noise ratio than the regular mammograms. The radiographs digitized with high resolution digitizers provide the morphological information of individual microcalcifications that makes the classification of microcalcifications into benign and malignant groups possible. The CNN system will need to be tested on regular mammograms. Some parameters of CNN may need to be fine tuned when applied to regular mammograms and the CNN system may not achieve the same performance level as it did in this study.

Both a large training and testing database are necessary in order to train and evaluate the performance of the neural network sufficiently and reliably. We will be expanding our database significantly in the future. To further improve the accuracy of the classification, a

hybrid neural network (HNN)⁶⁵ will also be employed to classify microcalcifications based on the input of both image data and image features⁶⁶ that will be automatically extracted.

The evaluation of feature based neural network with Generic Algorithm was based on a very difficult data set which was different from what was used for evaluation of image based neural networks. The microcalcification clusters in the data set for feature based neural network are very subtle and an experienced radiologist only scored an A_z of 0.54 in classifying between benign and malignant clusters. The size of the database is also small. We are currently in the process of expanding our database to include more mammograms from different hospitals.

Conclusions

We have demonstrated that the convolution neural networks can be an effective tool in the diagnosis of breast cancer. The results obtained in this study are very promising, even though they were based on a relatively small training and testing database. These results indicate the potential usefulness of CNN in classification of microcalcifications in digital mammograms. An extensive clinical test of our developed system using real mammograms will be needed to determine the clinical applicability.

Genetic algorithms are an effective tool to select optimal input features and structure of a backpropagation neural network. The neural network, combined with genetic algorithms, is able to effectively classify benign and malignant microcalcifications. The results of the neural network system can be used to help reducing the number of benign biopsies.

We have developed an image visualization system to aid radiologists to analyze breast MRI images efficiently and accurately. The interactive image visualization and quantitative image analysis enable radiologists to examine breast MRI cases more efficiently and accurately. We are also developing an automated detection system that can locate the potential tumors from the acquired images based on signal enhancement pattern and intensity. This visualization system can make BMRI more accurate and cost-effective, and thus make BMRI clinically feasible.

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Appendix

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